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MRI monitoring of lesions created at temperature below the boiling point and of lesions created above the boiling point using high intensity focused ultrasound*

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ABSTRACT

Magnetic Resonance Imaging (MRI) was utilized to monitor lesions created at temperature below the boiling point and lesions created at temperature above the boiling point using High Intensity Focused Ultrasound (HIFU) in freshly excised kidney, liver and brain and *in vivo* rabbit kidney and brain. T2-weighted fast spin echo (FSE) was proven as an excellent MRI sequence that can detect lesions with temperature above the boiling point in kidney. This advantage is attributed to the significant difference in signal intensity between the cavity and the thermal lesion. In liver the MRI sequence of Proton Density is recommended to detect lesions above boiling. In brain T1-W FSE was the optimum pulse sequence to detect lesions of either type. In order to monitor the temperature elevation during a HIFU exposure, T1-weighted fast spoiled gradient (FSPGR) was used. The shape of the focal temperature distribution was uniform with the absence of boiling, whereas with an exposure affected by boiling, the temperature distribution could be of irregular shape, demonstrating the drastic effects taking place during boiling. In order to confirm that boiling occurred, the temperature was estimated using the widely used method of Proton Resonance Frequency (PRF) shift.

Keywords: Ultrasound; Kidney; Brain; Liver; MRI; Lesion

1. INTRODUCTION

This paper is a continuation of previous papers by Damia-

nou [1] and Damianou *et al.* [2]. The first paper describes methods for ablating kidney tissue using high intensity focused ultrasound (HIFU), whereas the second paper deals with magnetic resonance imaging (MRI) guidance of HIFU for the case of kidney. This paper goes one step further by evaluating lesions created at temperatures above the boiling point during HIFU exposures using MRI. The evaluation was performed in kidney, liver and brain.

We have chosen to explore kidney, liver and brain because there is currently a lot of ongoing research either in animal models or in humans for these 3 tissues. In the area of kidney ablation with HIFU Watkin *et al.* [3] used a large animal model and proved the feasibility of this treatment method. Recently Roberts *et al.* [4] have performed ablations in the normal rabbit kidneys and they suggested that the mechanical effects of ultrasound, can be used to homogenize tissue.

HIFU ablation of renal tumours in humans remains in the early stages of clinical trials. In the early 1990s, Valancien *et al.* [5] reported the first clinical feasibility study in kidney using extracorporeal HIFU. Susani *et al.* [6], Wu *et al.* [7], and Marberger *et al.* [8] conducted clinical trials in patients with renal tumours and proved that HIFU may have a place in the treatment of renal tumours.

Hacker *et al.* [9] performed also ablation of 43 kidneys (porcine and human), using an experimental handheld extracorporeal technology. Finally Klingler *et al.* [10] use laparoscopic methods to treat kidney tumors.

Small animal models [11,12] were used to establish the feasibility of HIFU to create lesions in liver tissue. The thresholds for liver tissue destruction at varying exposure parameters were established in the 70s and 80s [13].

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Basic research in the area of liver continued during the 1990s; for example the histological effects of HIFU [14], the effects of blood perfusion on during liver ablation [15], and the relationship between tissue depth and the required intensity levels [16].

Several tumour models have been used to predict the effects of HIFU on liver tumors in humans (for example [17,18]) and to destroy VX2 liver tumours in rabbits [19, 20]. Hooded Sarcoma N (HSN) fibrosarcoma has been also used as a tumour model in rats with some success [21]. HIFU was also used for the treatment of metastatic melanoma in liver in a cat [22].

There is an increasing interest of work describing HIFU in the treatment of liver cancer in human clinical trials. In the early 1990s, Vallancien *et al.* [23] reported treatment of liver metastases using HIFU.

The Chongqing group published a study [24] describing a clinical trial of 68 patients with liver malignancies using HIFU. Li *et al.* [25] reported clinical trial of patients with liver cancer who were treated also with HIFU. Wu *et al.* [26] use HIFU in combination with TACE for the treatment of HCC.

MRI-guided HIFU has generally been used for the treatment of uterine fibroids using the ExAblate 2000 system (InSightec, Haifa, Israel). However, it is very likely that soon this system will be utilized for the treatment of kidney, liver and brain tumours. At Imperial College, London they have recently started a non-randomised clinical trial to assess the safety and efficacy of the MRI-guided HIFU system of InSightec in the treatment of liver tumours [27].

Thermal ablation of brain in animals with high intensity focused ultrasound (HIFU) was very popular in the 50's and 60's (for example [28] and [29]). HIFU was used in the clinical setting by Fry and Johnson [30] and showed that HIFU had the potential to treat brain cancer. Several groups used hyperthermia (heating of several minutes at 43°C) to treat brain tumours [31,32]. The clinical trials were abandoned probably due to the inexistence of effective imaging modality to guide the therapy. Especially for the case of brain it is extremely important to have absolute control of the ablation in order to avoid vital brain tissue damage such as the neurons. Now with the advancement of HIFU technology guided by MRI, it will be possible to conduct clinical studies for brain cancer.

In the early nineties several studies by the group of Dr. Hynynen [33-38] demonstrated the creation of lesions in animal brain and use MRI successfully for guiding and monitoring. Therefore, now there is increasing interest regarding brain ablation.

The combination of HIFU and MRI was first cited by

Jolesz and Jakab [39] who demonstrated that an ultrasonic transducer could be used inside a MRI scanner. The concept of using MRI to monitor the necrosis produced by HIFU was fully demonstrated in the early nineties in canine muscle [40-42]. In these studies it was shown that the contrast between necrotic tissue and normal tissue was excellent. This was a great enhancement for the HIFU systems because the therapeutic protocols can be accurately guided. Therefore the interest of using MRI as the diagnostic modality of guiding HIFU was increased. Although there are numerous studies regarding lesions created with temperature below the boiling point (also known as thermal lesions), there is insufficient information regarding MRI detection of lesions created with temperature above the boiling point. This paper focuses in the detection of lesions above boiling using MRI.

Several MRI sequences are investigated. For high quality imaging, this can be used at the end of a therapeutic protocol or at some instances of the protocol, the fast spin echo (FSE) techniques T1, and T2-weighted are investigated in all 3 tissues under investigation. For fast imaging, the T1-weighted fast spoiled gradient (FSPGR) MRI sequence was used. Fast imaging can be used to monitor the dynamic increase of tissue temperature during the application of an ultrasonic exposure. We have also use the fluid-attenuated inversion recovery (FLAIR) sequence for brain since this is the predominant sequence used in the clinical setting.

In order to prove whether a lesion is created at temperature below boiling or above, it is necessary to estimate the temperature at the focus. The proton resonance frequency (PRF) shift has been proven to be best pulse sequence for estimating temperature, because with this sequence the temperature is less dependent on the physiological changes of tissue during high-temperature HIFU exposures [43]. The temperature dependence of the PRF shift was measured to be linear above 50°C [43]. This linearity of the PRF shift above the tissue necrosis threshold allows the tissue temperature to be estimated during the therapeutic ultrasound exposures.

The task with high quality imaging was to find an optimum technique that can discriminate thermal from boiling lesions. Discriminating between lesion and normal tissue involves two types of tissues. Discriminating between thermal and boiling lesions involves three types of layers (normal tissue, lesion and cavity). Therefore the signal intensity vs. MR parameters needs to be evaluated for the above layers in order to optimise the contrast among tissue of interest, lesion and cavity. The other main task was to monitor the temperature elevation using a fast MRI technique in order to observe the shape

of the beam during HIFU exposures.

The growth of vapor bubbles due to boiling occurs due to the temperature induced by HIFU. Boiling is different from cavitation which occurs due to pressure oscillations induced by HIFU. Vapor bubbles created by boiling can grow rapidly to a size of few millimetres [44]. This growth can be explosive due to the super heating caused by HIFU. Therefore the cavities produced of few mms can be easily monitored by MRI.

There are two reasons for studying the MRI appearance of boiling lesions that could be of importance:

1) Since boiling provides enhanced heating, it could be possible for large tumours (for example giant fibroadenomas) to use this type of heating (especially in the center of the tumor) in order to accelerate the ablation,

2) since the rabbit model is used extensively by many researchers in MRI animal experiments (for example [33-37]), then it is very useful for them to know the appearance of boiling lesions.

2. MATERIALS AND METHODS

2.1. HIFU/MRI System

Figure 1 shows the block diagram of the HIFU/MRI system which includes the following subsystems:

1) HIFU system, 2) MR imaging, 3) Positioning device (robot) and associate drivers, 4) Temperature measurement, 5) Cavitation detection, 6) MRI compatible camera, 7) Software.

2.1.1. HIFU System

The HIFU system consists of a signal generator (HP 33120A, Agilent technologies, Englewood, CO, USA), a RF amplifier (250 W, AR, Souderton, PA, USA), and a spherically shaped bowl transducer made from piezoelectric ceramic of low magnetic susceptibility (Etalon, Lebanon, IN, USA). The transducer used for the kidney and liver ablation operates with frequency of 4 MHz, and the transducer for brain ablation operates at 1 MHz. The transducer is rigidly mounted on the MRI-compatible positioning system (MEDSONIC LTD, Limassol, Cyprus) which is described shortly.

2.1.2. MRI Imaging

The 3-d positioning device and the transducer were placed inside a MRI scanner (Signa 1.5 T, by General Electric, Fairfield, CT, USA). The spinal coil (USA instruments, Cleveland, OH, USA) was used to acquire the MRI signal for the case of kidney and liver, whereas a brain coil (USA instruments, Cleveland, OH, USA) was used to acquire the MRI signal from the brain tissue.

2.1.3. Positioning Device/Drivers

The robot has been developed initially for three de-

grees-of-freedom, but it can be easily developed for 5 degrees of motion. Since the positioning device is placed on the table of the MRI scanner its height should be around 55 cm (bore diameter of the MRI scanner). The length of the positioning device is 45 cm and its width 30 cm. The weight of the positioning device is only 6 kg and therefore it can be considered portable. **Figure 2** shows the schematic the positioning device illustrating the 3 stages, transducer, and coupling method. The positioning device operates by means of 3 piezoelectric motors (USR60-S3N, Shinsei Kogyo Corp., Tokyo, Japan). More details of this positioning device can be found in [31]. The movement of the positioning device is monitored by an MRI compatible camera (not shown in the figure) which is placed on a non-magnetic holder. Moreover, the positioning system includes optoelectronic encoders for providing signals indicating the relative positions of the movable elements in the positioning system. The resolution of all 3 axes of the positioning device is 0.1 mm.

2.1.4. MRI Compatible Camera

In order to monitor the condition of the animal or humans

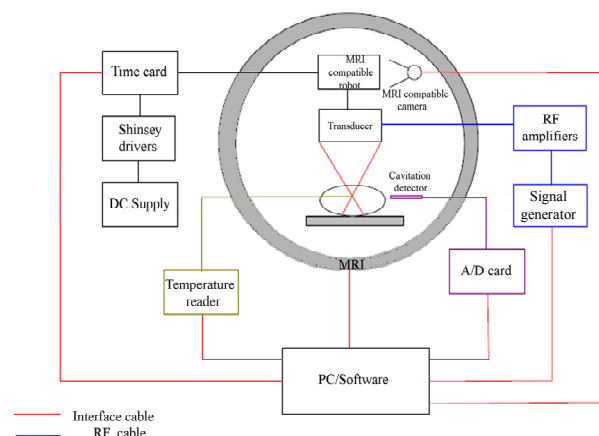


Figure 1. HIFU system under MRI guidance showing the various functionalities of the HIFU/MRI system.

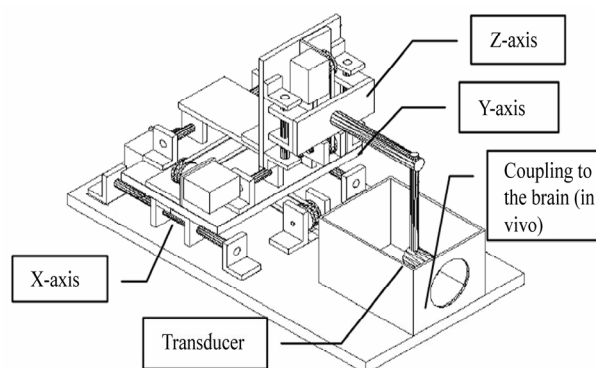


Figure 2. Schematic of the robot showing all of its stages.

(future use), a MRI compatible camera (MRC Systems GmbH, Heidelberg, Germany) was mounted on the system. The camera was interfaced by means of a video card. With the aid of the MRI compatible camera, the researcher can monitor the welfare of the animal.

2.1.5. Temperature Measurement

Temperature was measured in few experiments in order to confirm that the temperature estimated using the PRF method was accurate enough. Temperature is measured using a data acquisition system (HP 34970A, Agilent technologies, Englewood, CO, USA). Temperature is sensed using a 50- μ m diameter T-type copper-costantan thermocouple (Physitemp Instruments, Inc. New Jersey, USA) which is MRI compatible. The thermocouple is placed in the tissue by means of a catheter. The thermocouple measures the temperature at the focus. This is achieved by applying low-intensity (low enough not to cause tissue damage) and during the application of ultrasound the transducer is scanned accordingly in order to detect the maximum temperature. This establishes positioning of the thermocouple in the focus of the transducer. The temperature error of the thermocouple is in the order of 0.1°C.

2.1.6. Cavitation Detector

From a scientific point of view it will be useful to separate lesions developed based on thermal and based on cavitation mechanisms. In this system we use a passive MRI compatible cavitation detector (Etalon, Lebanon, IN, USA), which is placed perpendicularly to the beam of the HIFU transducer (method described in [45]). Since the HIFU protocol is applied inside the magnet of an MRI scanner, the detector must be MRI compatible. The diameter of the detector was 1 cm, its radius of curvature was 10 cm and operated between 1 and 13 MHz (centre frequency is 7 MHz). The detector was mechanically coupled to the HIFU transducer. The voltage from the detector was fed to an analogue to digital (A/D) card (CS1250, A/D 12 bit, 50 MHz, from GAGE, Lachine, Canada). The A/D card was synchronised to receive the signal when the HIFU transducer was activated. The received signal was stored in a PC (Dell Inc. Round Rock, Texas, USA). The signal was then displayed using EXCEL (Microsoft Corporation, Redmond, WA USA) in order to visualize whether cavitation occurs or not.

2.1.7. Software

A user-friendly program written in MATLAB (The Mathworks Inc., Natick, MA) has been developed in order to control the system. The software serves the following main tasks: 1) Displaying of MR images, 2) transducer movement (the user may move the robotic arm in a specific direction or customize the automatic movement of the robotic arm in any rectangular forma-

tion by specifying the pattern, the step and the number of steps), 3) messaging (starting time, treatment time left etc), 4) Patient data (age, weight, etc), 5) Display of motor position, and 6) Display of the contents of an MR compatible camera, 7) Cavitation detection window, 8) temperature measurement, and 9) MR estimated temperature using the PRF method.

2.2. In Vitro Experiments

The tissue was placed on top of an absorbing material in order to shield adjacent tissue from stray radiation from the bottom. The transducer was placed on the arm of the positioning device and was immersed in the water tank, thus providing good acoustical coupling between tissue and transducer. Any bubbles that may have collected under the face of the transducer face were removed in order to eliminate any reflections. In all experiments the tissues used (kidney, liver and brain) were extracted from freshly killed lamb, and the experiment was conducted in the same day. Totally 22 kidneys, 8 liver and 16 brains were ablated for investigating various issues.

2.3. In Vivo Experiments

For the *in vivo* experiments, adult rabbits from Cyprus were used weighting approximately 3.5-4 kg. Totally 8 rabbits were used in the experiments. The rabbits were anaesthetized using a mixture of 500 mg of ketamine (100 mg/mL, Aveco, Ford Dodge, IA), 160 mg of xylazine (20 mg/mL, Loyd Laboratories, Shenandoah, IA), and 20 mg of acepromazine (10 mg/mL, Aveco, Ford Dodge, IA) at a dose of 1 mL/kg.

Presence of the skull in the ultrasonic path not only distorts the field by reflection, but may also destroys the underlying tissue in contact with it by absorbing ultrasonic energy and dissipating it as heat. A craniotomy adequate in extent to permit unimpeded passage of the cone of sound was imperative. The extent of the craniotomy depends on the solid angle of radiation and the depth of the target from the cranial surface. The larger the angle and the deeper the target, the larger the size of craniotomy needed. For the transducer used and a target depth of 1 cm, a circular craniotomy of 3 cm in diameter was adequate. The animal experiments protocol was approved by the national body in Cyprus responsible for animal studies (Ministry of Agriculture, Animal Services).

2.4. HIFU Parameters

The *in situ* spatial average intensity was estimated based on the applied power and the half-power width of the beam of the transducer. The details of the intensity estimation can be found in Damianou 2003 [1]. In order to create large lesions, a grid pattern of 3 \times 3 or 4 \times 4 over-

lapping lesions was used. The spacing between successive transducer movements was 2 mm, which creates overlapping lesions for the intensity and pulse duration used. In all the exposure, unless stated otherwise the ultrasound was turn on for 5 s. The delay between successive ultrasound firings was 10 s for the scanned ablation. The intensity indicated through this paper is in situ spatial average intensity.

2.5. MRI Parameters

The various MRI parameters used for the various pulse sequences are listed in **Table 1**. The Region of Interest (ROI) was circular with diameter of nearly 2 mm.

The temperature change (T) has been estimated using the equation stated in Chung *et al.* 1996 [46] which is as follows:

$$\Delta\phi = \gamma B_0 \alpha \Delta T T E \quad (1)$$

where $\Delta\phi$ is the temperature-dependent phase shift which is the phase acquired before and during temperature elevation and which accumulates during the echo time TE using the gradient-echo pulse sequence FSPGR. The other terms are γ which is the gyromagnetic ratio of proton, 42.58 MHz/T, α is the average proton resonance frequency coefficient and B_0 is the flux density of the static magnetic field. The measured temperature elevation can be added to the base-line temperature to obtain the absolute temperature. The average temperature coefficient for the frequency shift was taken from the study of Vykhodtseva *et al.* [47].

3. RESULTS

Figure 3 shows an MRI image of 4 lesions *in vitro* kidney (plane perpendicular to the beam) resulting from intensities ranging from 1000 to 2500 W/cm² using T1-weighted FSE (**Figure 3(a)**), T2-weighted FSE (**Figure 3(b)**) and T1-weighted FSPGR (**Figure 3(c)**) with pulse duration of 5 s. The MRI parameters used are shown in **Table 1** (row 1, 2 and 4). The MRI estimated maximum temperature at the focus was 55°C for the intensity of 1000 W/cm², 83°C for 1500 W/cm², 105°C for 2000 W/cm² and 123°C for 2500 W/cm². The

temperature measured using the thermocouple for the 1000 W/cm² was 53°C, which is very close to the temperature estimated using the PRF method.

Note that with T2-weighted FSE (**Figure 3(b)**) white spots (cavity) within the dark thermal lesion are seen for intensities higher than 2000 W/cm². T1-W FSE and FSPGR show some indication of these cavities, but the resolution is weaker than T2-w FSE.

Figure 4 shows MRI images (in a plane parallel to the transducer beam) of 3 lesions in kidney *in vitro* at intensities of 1000, 2000 and 3000 W/cm². Again T2-weighted FSE shows cavities (due to boiling) within the thermal lesion. T1-W FSE and T1-W FSPGR fail to provide good resolution in this axis.

Having observed that T2-W FSE was probably a successful MRI sequence to detect boiling lesions, this pulse sequence was investigated further by evaluating the Contrast to Noise Ratio (CNR) vs. Echo Time (TE). **Figure 5** shows the plot of CNR vs. TE for kidney tissue, lesion and cavity of the lesion of **Figure 4(b)** (intensity of 3000 W/cm²) demonstrating that good contrast between lesion and cavity is achieved using T2-weighted FSE between 20 and 50 ms.

Figure 6 shows a T2-weighted FSE image of 3 lesions in rabbit kidney *in vivo* using different intensities (1000, 2000 and 2500 W/cm²) for a 5 s pulse. The estimated temperature was 52°C, 95°C and 115°C.

The lesion created using 2500 W/cm² appears to have a white spot (cavity). These temperatures are lower than the corresponding temperatures *in vitro* (**Figure 4**) for the same intensity due to the removal of heat due to blood flow or possibly due to reflection from various interfaces.

Figure 7(a) shows an MR image of a lesion acquired using T1-weighted FSPGR (for MRI parameters see **Table 1**, row 4). **Figure 7(b)** shows the photograph of the kidney showing cavity within the large thermal lesion.

Note that this large lesion was created using a matrix of 4 × 4 single lesions with intensity of 1500 W/cm². Out of these 16 lesions, one lesion was created possibly due to the mechanism of cavitation, which results to tissue evaporation or boiling. The estimated temperature during

Table 1. Parameters used for the various MRI pulse sequences.

Series	NAME	TR (ms)	TE (ms)	Slice thickness (mm)	Matrix	FOV (cm)	NEX	BW (KHz)	ETL	Other
1	T1-weighted FSE	500	9.2	3, (gap 0.3 mm)	256 × 256	16	1	31.25	8	-
2	T2-weighted FSE	2500	8,16,32,48,64,80	3, (gap 0.3 mm)	256 × 256	16	1	31.25	8	-
3	PD	2500	7.2	3, (gap 0.3 mm)	256 × 256	16	1	31.25	8	-
4	FSPGR T1-weighted	50	2.7	3, (gap 0.3 mm)	256 × 256	16	1	62.50	-	Flip angle 50
5	FLAIR	8000	80	3, (gap 0.3 mm)	256 × 256	16	1	6.9	8	Inversion Time 1200-2200

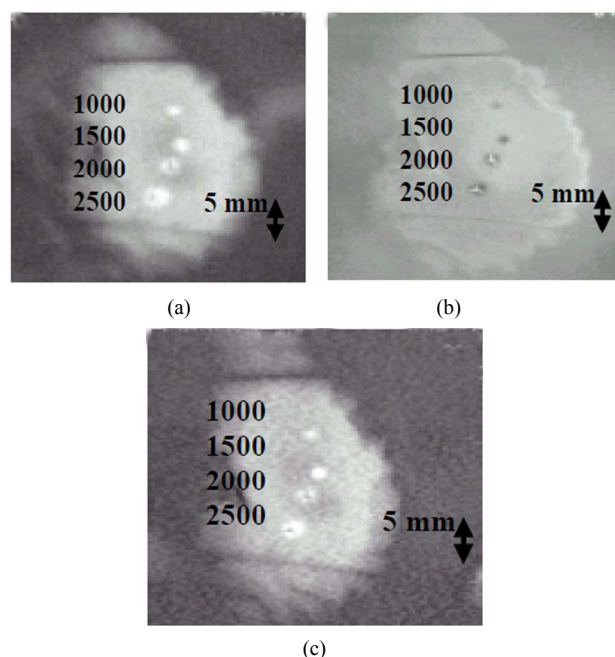


Figure 3. MR images (in a plane perpendicular to the beam) of four lesions (intensities 1000, 1500, 2000 and 2500 W/cm²) in kidney *in vitro* using. (a) T1-weighted FSE; (b) T2-weighted FSE; (c) T1-weighted FSPGR. With intensities above 2000 W/cm² the lesions exhibit boiling activity. The discrimination between boiling and non-boiling lesion is best monitored using T2-W FSE.

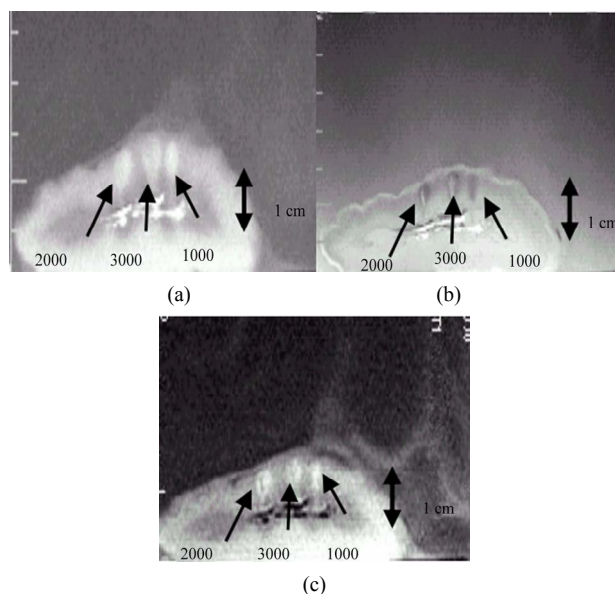


Figure 4. MR images (in a plane parallel to the beam) of 3 lesions in kidney *in vitro* using. (a) T1-weighted FSE; (b) T2-weighted FSE; (c) T1-weighted FSPGR. With intensities above 2000 W/cm² the lesions exhibit boiling activity. The discrimination between boiling and thermal lesion is best monitored using T2-W FSE.

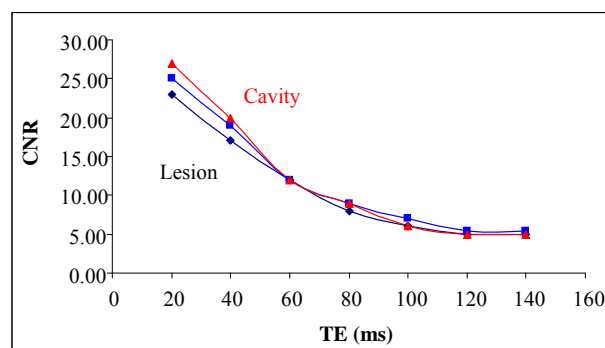


Figure 5. CNR vs. TE for the lesion, kidney and cavity for the lesion of **Figure 8(b)** with intensity of 3000 W/cm².

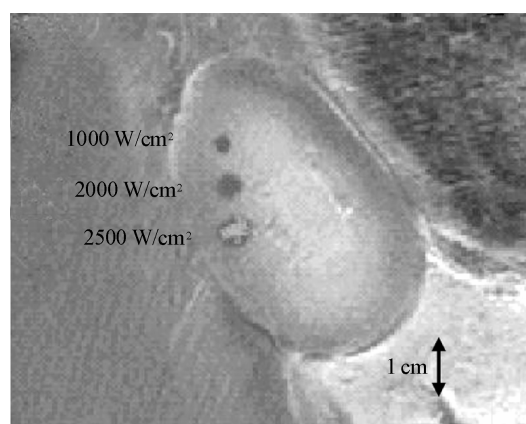


Figure 6. MRI image using T2-weighted FSE of 3 lesions in rabbit kidney *in vivo* at different intensities (1000, 2000, and 2500 W/cm²) for a 5 s pulse. The lesion with intensity of 2500 W/cm² is affected by tissue boiling.

the creation of this lesion was 120°C, whereas the temperature for the rest of the lesions (non-boiling) varied from 80 to 85°C.

This bubbly lesion exhibits low-signal lesion (dark spot) and lies inside the large lesion (white spot within the kidney tissue). At the location of the boiling lesion, the passive cavitation detector confirmed the occurrence of cavitation since broadband emission was detected (see **Figure 7(c)** which shows the frequency spectrum of the HIFU transducer). Although with this intensity bubbly lesions should not be produced, the high temperature estimated for this one lesion, should be attributed to cavitation. Cavitation was possibly initiated by bubbles that are trapped in the *in vitro* tissue due to the absence of blood in the vasculature.

Figure 8 demonstrates the temperature increase *in vitro* kidney using HIFU and MRI monitoring. The MRI images were acquired using the dynamic sequence T1-weighted FSPGR. Each image was acquired in 5 s. In **Figure 8(a)** ultrasound was OFF. In the next 5 images

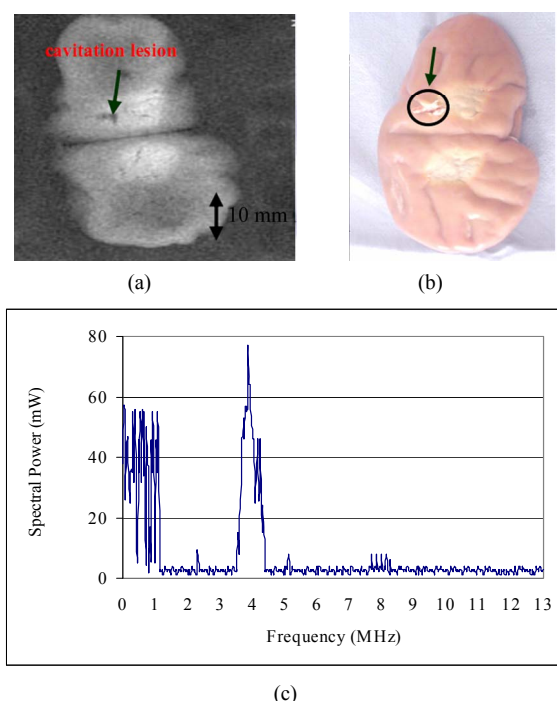


Figure 7. (a) MR image (in a plane perpendicular to the beam) of large lesion in kidney *in vitro* using T1-weighted FSPGR (TR = 50 ms), showing one cavitation lesion; (b) Photograph of the kidney showing the cavitation lesion within the large thermal lesion; (c) Frequency spectrum of the HIFU transducer exhibiting cavitation activity.

the applied spatial average intensity was 1000 W/cm^2 (for 25 s), and in the last 2 images ultrasound is turned OFF. Due to the heating a dark spot is observed (see arrows). The estimated maximum temperature was 55°C .

Figure 9 shows the corresponding temperature increase using T1-weighted FSPGR influenced by boiling. In **Figure 9(a)** ultrasound was OFF. In the next 5 images the applied spatial average intensity was 3500 W/cm^2 (for 25 s). Note that compared to **Figure 7** where the focal beam is circular, the focal beam in this figure is distorted, which is attributed to the occurrence of boiling confirmed also by the temperature of 112°C measured.

Figure 10 shows MRI image of lesion in liver. The image was acquired using Proton density. The lesion was created using low intensity (1000 W/cm^2) for long time (30 s). These ultrasonic parameters produce temperature elevation which is above the boiling point (120°C). Thus, the evaporation of tissue causes a cavity that follows the shape of the beam.

Figure 11 shows the MRI image a large lesion in lamb brain *in vitro* created by scanning the transducer with a 4×4 grid using 2000 W/cm^2 using T1-w FSE resulting to a large bubbly lesion. The maximum estimated temperature for this lesion was 110°C . Note that

in some location no lesion was created due to poor ultrasound penetration due to the air bubbles possibly trapped in the blood vessels. This image shows once more the excellent contrast between normal brain and lesions (in this case boiling lesions).

Figure 12 shows ablation in rabbit *in vivo* using a 4×4 grid with intensity of 2000 W/cm^2 for 20 s. This large lesion was created using thermal mechanisms and therefore the lesion appears bright. The maximum estimated temperature for this exposure was 90°C . The contrast of thermal lesions is definitely much better than the case of boiling lesions. Unlike the *in vitro* case of **Figure 11** where with this level of intensity boiling lesions were created, in this *in vivo* example in none of the 16 lesions a boiling lesion was produced. This proves that *in vitro* brain includes bubbles which are responsible for producing boiling lesions possibly due to enhanced heating or cavitation. Both of these mechanisms possibly produced temperatures above boiling and therefore bubbly lesions were created.

The methodology applied for kidney which evaluates T1-W FSE, T2-W FSE and FSPGR (**Figures 3 and 5**) was also applied for liver. In brain in addition to these 3 pulse sequences, FLAIR was also evaluated because this MRI sequence is widely used clinically for the case of brain. It was found that for liver the best pulse sequence to evaluate boiling lesions was T2-w FSE with minimum TE (*i.e.* proton density). For brain T1-W FSE was the best pulse sequence to monitor both boiling and non-boiling lesions. **Table 2** summarizes the recommended MRI sequence according to our experience for monitoring boiling lesions.

4. DISCUSSION

So far it was concluded by several studies (for example Quesson *et al.* [48], Salomir *et al.* [49], McDannold *et al.* [50]) that MRI-guidance of HIFU serves mainly 4 purposes: 1) localisation of the focus, 2) imaging of the temperature elevation, 3) imaging at the end of the treatment protocol, in order to evaluate the necrosis and

Table 2. Recommended pulse sequences for discriminating between (a) normal and thermal lesions; and (b) normal tissue, lesions created with temperature below boiling and lesions created with temperature above boiling.

Tissue type	Contrast between lesion below boiling and tissue	Contrast between lesion below boiling, tissue, and lesion above boiling
Kidney	T1-W FSE, T2-W FSE	T2-W FSE
Liver	T1-W FSE, T2-W FSE	T2-W FSE
Brain	T1-W FSE	T1-W FSE

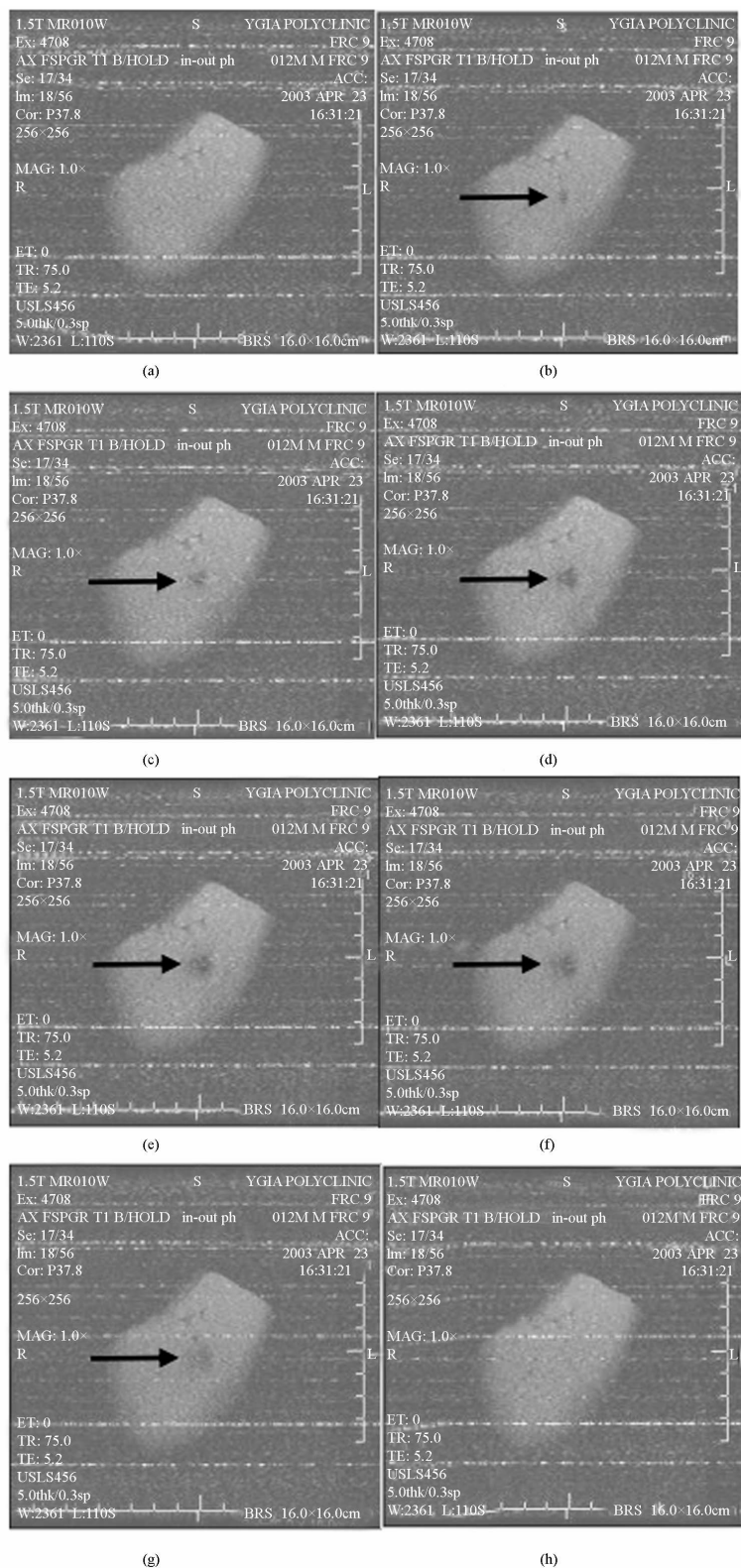


Figure 8. Temperature evolution *in vitro* kidney using T1- weighted FSPGR (thermal mechanism). Each image was acquired in 5 s. (a) Ultrasound is OFF; (b)-(f) Applied spatial average intensity: 1000 W/cm² (for 25 s); (g)-(h) Ultrasound is OFF.

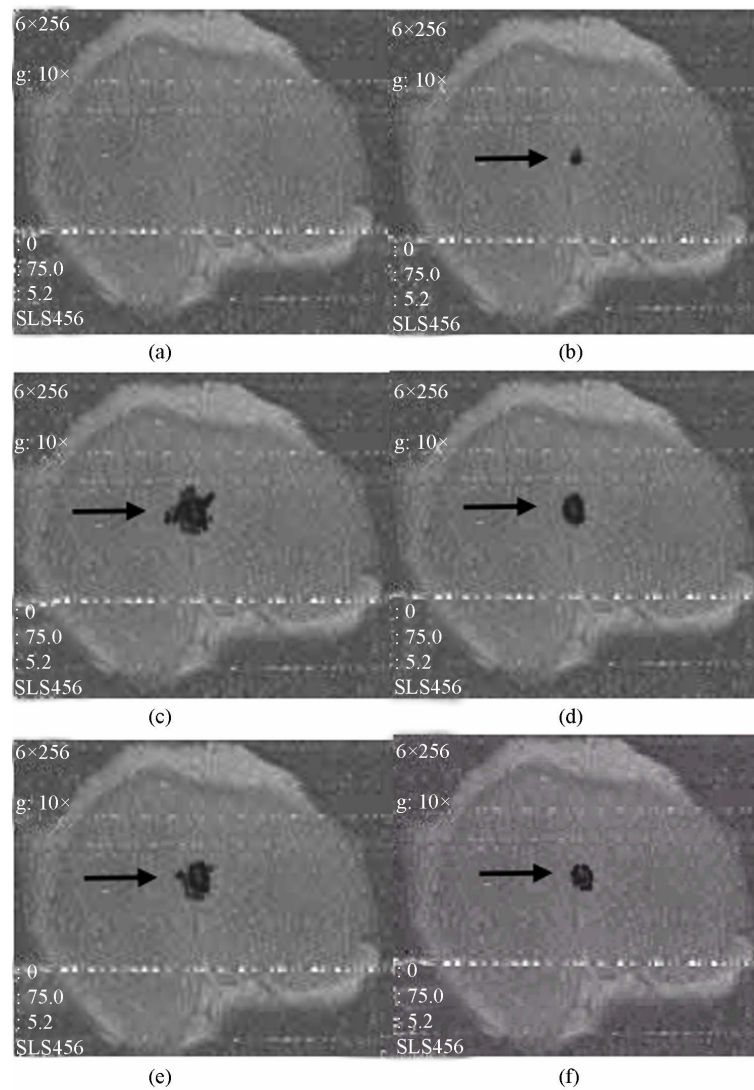


Figure 9. Temperature elevation *in vitro* kidney using T1- weighted FSPGR (boiling). (a) Ultrasound is OFF; (b)-(f) applied spatial average intensity: 3500 W/cm^2 (for 25 s).

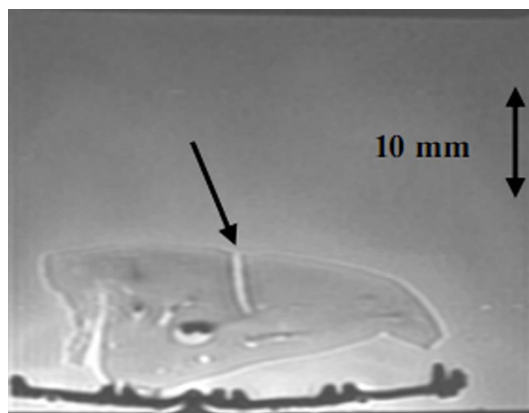


Figure 10. MR image using proton density of lesion in liver created under the influence of boiling. The lesion was created using low intensity (1000 W/cm^2) for long time (30 s).

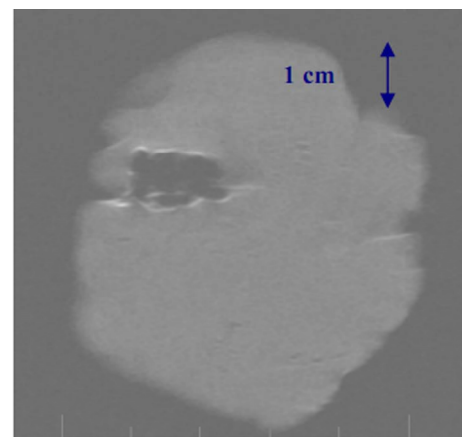


Figure 11. MRI image using T1-w FSE of large boiling lesion created *in vitro* using 2000 W/cm^2 for 20 s.

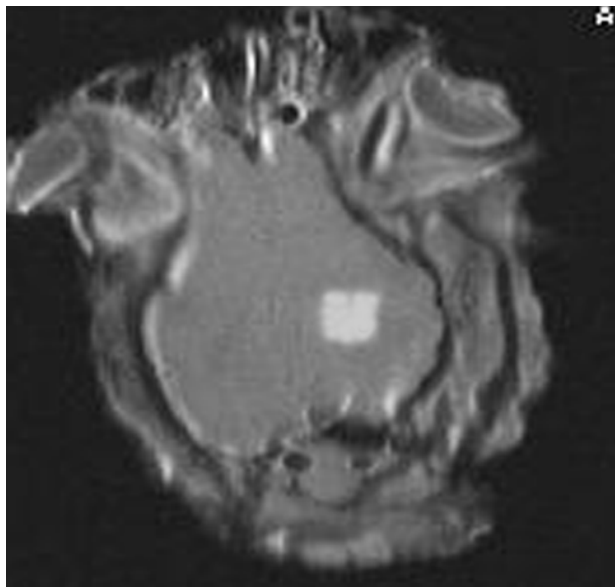


Figure 12. MRI image using T1-w FSE of large thermal lesion created *in vivo* using 2000 W/cm² for 20 s.

possibly re-planning the treatment protocol in the event of incomplete coverage of the target and 4) follow-up imaging, evaluating the effectiveness of HIFU ablation, several days after the treatment. The main goal in this paper was to develop methods for detecting lesions created at temperature below boiling point and lesions created above boiling using MRI techniques.

This paper enhances the role of MRI guidance in HIFU because it provides techniques to discriminate between non-boiling and boiling lesions.

In kidney the best MRI sequence to detect boiling lesions was T2-W FSE. T2-W FSE was evaluated further by plotting the CNR vs. TE for the three regions of interest (kidney, lesion and cavity). It was concluded that between TE's of 20 and 50 ms, the signal difference, and hence the contrast between the three regions of interest (kidney, lesion and cavity) is maximum (**Figure 5**). According to **Figure 5**, the cavity, which appears inside the lesion, has stronger signal and decays slower.

T1-weighted FSE and T1-weighted FSPGR do not consistently show boiling lesions in kidney, and even when cavities are visible the contrast is not very good.

In liver T2-W FSE with low TE (*i.e.* Proton density) is recommended as an MRI sequence to detect non-boiling lesions and boiling lesions.

Previous literature [36,37] demonstrated that lesions in the brain can be monitored with excellent contrast using T1-W FSE. However in the previous studies only thermal lesions were shown. In this paper we have explored extensively the use of MRI to image both lesions created under thermal mechanisms and under boiling.

With T1-w FSE the signal intensity of the brain tissue is homogeneous and therefore the contrast with thermal lesions or with bubbly lesions is excellent. In this study it was concluded that in brain T1-weighted FSE was the optimum MRI sequence not only to detect non-boiling lesions, but also boiling lesions.

It was observed that bubbly lesions appear darker than thermal lesions. Bubbly lesions appear dark, due to the air spaces resulting from cavities caused by boiling or sometimes by cavitation. In the brain tissue *in vitro* it was demonstrated that non-degassed excised tissue is a good model for easily initiating enhanced heating or cavitation since air is trapped in the vasculature of the brain. This model of cavitation might not be of any significant for clinical use since in live tissue cavitation will not occur with such exposure; however, this model of initiating cavitation is very useful for the purpose of studying the MRI appearance of bubbly lesions. The exposure of using 2000 W/cm² in brain results to temperatures of around 90°C (**Figure 12**) *in vivo*. However with the same exposure *in vitro* the resulting estimated maximum temperature was 110°C. This is probably due to enhanced heating or cavitation which results to tissue boiling. The enhanced heating due to bubbles was also noted in the studies by Fry *et al.* [51], by ter Haar [52], and by Chavier *et al.* [53].

This is the first paper demonstrating creation of large lesions in brain formed by scanning the transducer in grid formation. Both thermal and bubbly lesions were monitored successfully using T1-w FSE with excellent contrast, proving the potential of HIFU to treat reliably tumours in the brain in the future. The proposed system effectively creates large lesion in brain and on the same time these lesions are effectively monitored using MRI with strong confidence on the margins of these lesions especially using T1-w FSE.

Boiling bubbles scatter and reflect ultrasound. These reflections result in shielding the HIFU focus and thus increased prefocal heating is observed. Therefore boiling bubbles, similar to cavitation bubbles, result to distortion of the lesion from a cigar shape into a tadpole shape (**Figure 4**). In addition to this the growth of the lesion is shifted towards the transducer. This phenomenon which is attributed to bubbles, either due to ultrasound-induced cavitation or to boiling has been observed also in the study by ter Haar [53], and by Chavier *et al.* [52]. For the scanned lesions the focus is not shifted towards the transducer, because while boiling begins in a very short time in both single and scanned lesions, it cannot distort the lesion for scanned exposures because the focus of the transducer moves away from the boiling site. This speculation is also supported in the paper by Khokhlova [44].

This paper also showed by means of MRI images that focal beam is distorted during the occurrence of boiling. Initially low intensity was used (1000 W/cm^2) and the temperature elevation was monitored using T1-weighted FSPGR. The shortest acquisition time that we could achieve with our system was 5 s. Modifying any MRI parameter to decrease the time resulted to poor contrast or low signal to noise ratio (SNR). If we had achieved faster acquisitions (less than 5 s), then it would have possible to see more drastic changes. A decrease in the signal (black spot) demonstrated the increase in the temperature (also observed by Hynynen *et al.* [54]). The shape of the black spot was circular. The black spot increased gradually with increased temperature, and the shape remains circular at all times (**Figure 8**). When the intensity was increased to 3500 W/cm^2 , the shape of the black spot was irregular indicating that boiling occurred (**Figure 9**).

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